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**2) Over-expression of the MDM2 gene is found in some cases of haematological malignancies .**

Quesnel B; Preudhomme C; Oscier D; Lepelley P; Collyn-d'Hooghe M; Facon T ; Zandecki M; Fenaux P

Inserm U124, Institut de Recherches sur le Cancer de Lille, France.

British journal of haematology (ENGLAND) Oct 1994 , 88 (2) p415-8,  
ISSN 0007-1048 Journal Code: 0372544

**3) Amplification of the MDM2 gene in human breast cancer and its association with MDM2 and p53 protein status.**

McCann A H; Kirley A; Carney D N; Corbally N; Magee H M; Keating G; Dervan P A

Biotechnology Centre, University College Dublin, Belfield, Ireland.

British journal of cancer (SCOTLAND) May 1995 , 71 (5) p981-5,  
ISSN 0007-0920 Journal Code: 0370635

**4) Frequent occurrence of p53 mutations in rhabdomyosarcoma and leiomyosarcoma, but not in fibrosarcoma and malignant neural tumors.**

Wurl P; Taubert H; Bache M; Kroll J; Meye A; Berger D; Siermann A; Holzhausen H J; Hinze R; Schmidt H; Rath F W

Surgical Clinic, Martin Luther University of Halle-Wittenberg, Halle/S., Germany.

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## FREQUENT OCCURRENCE OF *p53* MUTATIONS IN RHABDOMYOSARCOMA AND LEIOMYOSARCOMA, BUT NOT IN FIBROSARCOMA AND MALIGNANT NEURAL TUMORS

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We have analyzed soft-tissue sarcomas (STS) molecularly for mutations in the tumor-suppressor gene *p53* and immunohistochemically for expression of *p53* and *mdm2* proteins. In this study, tumor samples from 3 groups of soft-tissue sarcomas, i.e., fibrosarcomas, myogenic sarcomas and malignant neural tumors (MNT), were investigated. The methods applied encompass immunohistochemistry on 198 tumor samples using *p53* antibodies (DO-1 and DO-7) and an *mdm2* antibody (IF-2). Out of these, 100 samples were subjected to non-radioactive PCR-SSCP-sequencing analysis. Immunohistochemical detection rate for *p53* (range of 57% to 67%) and for *mdm2* proteins (range of 19 to 44%) was similar in all 3 groups. In higher tumor grades, an increased rate of immunopositivity was found for *p53* but not for *mdm2*. Investigation of *p53* mutational status revealed 6 mutations in myogenic sarcomas but none in malignant neural tumors or fibrosarcomas, suggesting different roles of *p53* in the 3 STS groups. Interestingly, a G → A transition in codon 245 (a CpG site) was found in 3 myogenic sarcomas. Our results and those of others suggest *p53* codon 245 as a mutational hotspot in sarcomas, as recognized in carcinomas.

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Soft-tissue sarcomas (STS) can be defined as malignant tumors of non-epithelial extraskelatal tissue, excluding the reticulo-endothelial system, glia, and supporting tissue of various parenchymal organs. By convention, malignant tumors of the peripheral nervous system are included (Enzinger *et al.*, 1969). In addition to MFH's and liposarcomas, the most frequent are fibrosarcomas (8–12%), malignant peripheral-nerve-sheath tumors (8–10%), peripheric neuroblastomas (5%), leiomyosarcomas (5–10%), and rhabdomyosarcomas (10–20%) (Enzinger *et al.*, 1969; Enjoji and Hashimoto, 1984). However, even after distinct tumor classification, there are no comprehensive immunohistochemical and molecular data that convincingly characterize malignant tumors according to the course of disease and the prognosis. Apart from oncogenes, tumor-suppressor genes, in particular, and their role in cell-cycle regulation are of crucial interest. Among tumor suppressors, *p53* stands out, with about 50% of mutational alterations in malignomas (Hollstein *et al.*, 1991, 1996). *p53* mutational analysis for soft-tissue sarcomas has been performed for fibrosarcomas (Latres *et al.*, 1994), neuroblastomas (Imamura *et al.*, 1993; Komuro *et al.*, 1993; Vogan *et al.*, 1993; Hosoi *et al.*, 1994), neurofibrosarcomas (Menon *et al.*, 1990), leiomyosarcomas (Stratton *et al.*, 1990; Andreassen *et al.*, 1993; Liu *et al.*, 1994; Latres *et al.*, 1994; Patterson *et al.*, 1994; Cordon-Cardo *et al.*, 1994; De Vos *et al.*, 1994), rhabdomyosarcomas (Stratton *et al.*, 1990; Cordon-Cardo *et al.*, 1994; Mulligan *et al.*, 1990; Castresana *et al.*, 1995; Felix *et al.*, 1992), MFH's and liposarcomas (reviewed in Taubert *et al.*, 1995), and other STS (Liu *et al.*, 1994; Cordon-Cardo *et al.*, 1994; Toguchida *et al.*, 1992; Boman *et al.*, 1994; Scinicariello *et al.*, 1994; Dumaz *et al.*, 1993; Hollstein *et al.*, 1994). In carcinomas, the great majority of *p53* mutations are missense mutations, and out of these about 40% occur at mutational hot spots (Levine, 1993). On closer examination, one third of 280 tumor mutations was found to consist of transitions at hot-spot regions with CpG sites (Hollstein *et al.*, 1991). On investigating the mutational spectrum for soft-tissue sarcomas, we found a spectrum similar to that of carcinomas (Taubert *et al.*, 1995). Most of the muta-

tions are missense mutations, with the majority occurring at CpG sites.

In addition to mutations in the *p53* gene, an amplification of the *mdm2* oncogene affects sarcoma tumorigenesis. Amplification of the *mdm2* gene results in *mdm2* protein over-expression with complexing and inactivating *p53* protein (Momand *et al.*, 1992). It was detected in liposarcomas, MFH's (Oliner *et al.*, 1992; Leach *et al.*, 1993), leiomyosarcomas (Patterson *et al.*, 1994), a group of different soft-tissue sarcomas (Cordon-Cardo *et al.*, 1994) and osteosarcomas (Oliner *et al.*, 1992; Ladanyi *et al.*, 1993). Immunohistochemical detection of *mdm2* over-expression revealed that in most cases positive staining is alternative to *p53* alterations over-expression (Leach *et al.*, 1993), but we found co-existing over-expression, earlier described for soft-tissue sarcomas (Cordon-Cardo *et al.*, 1994).

The goal of our study was to extend the mutational analysis for *p53* on 3 sarcoma entities with high occurrence: myogenic sarcoma (Leiomyosarcoma and rhabdomyosarcoma), fibrosarcoma and malignant neural tumors. We also examined these tumors for alterations in *K-ras* and *N-ras* genes, in order to more fully comprehend the mutational spectrum in malignant soft-tissue tumors. Additionally, we tested different monoclonal antibodies (MAbs) against *p53* and *mdm2* for a large number of patients and tumor specimens. The immunohistochemical and mutational data were considered in relation to clinical data, to improve understanding of their clinical relevance.

### MATERIAL AND METHODS

#### Tumor samples

A collection of 198 tumor samples originating from formalin-fixed paraffin-embedded STS from 127 patients (Institute of Pathology and Surgical Clinic, University of Halle, Germany) were chosen for immunohistochemistry (IHC). Of these, 100 samples from 78 patients were investigated for *p53* mutations by PCR-SSCP-sequencing analysis (Table I). For *K-ras* mutation, we investigated 32 samples from myogenic sarcomas (21 patients) and, for detection of *N-ras* mutation, 31 MNT samples (25 patients) were examined. All the patients had had local radical surgical treatment. From the patients involved, clinical data, including the survival rate, were collected and grading was performed, taking into consideration mitotic activity and verification of tumor necrosis (Van Unnik *et al.*, 1988). Patients average post-treatment observation periods were 47 months (7 to 130) for fibrosarcomas, 22 months (2 to 107) for malignant neural tumors, 30 months (1 to 156) for leiomyosarcomas and 17 months (7 to 30) for rhabdomyosarcomas.

#### Immunohistochemistry

Immunohistochemical staining for *p53* was done for all tumor samples, using MAbs DO-1 (Oncogene Science, Manhasset, NY) and DO-7 (Medac, Hamburg, Germany). Samples

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with *p53* mutations were characterized additionally by a polyclonal antibody CM-1 (Medac) and 2 other MAbs Pab 1801, Pab 240 (Oncogene Science) as described (Taubert *et al.*, 1995). For mdm2 we used the MAb IF2 (Oncogene Science) recognizing an N-terminal epitope. IF-2 was used at a working solution of 5 µg/ml. All other steps of staining were the same as described (Taubert *et al.*, 1995). As positive control for mdm2 the osteosarcoma cell line Saos-2 and for *p53* Panc TuI were used. As negative control, the first antibody was omitted and replaced by an unrelated MAb of the same isotype in the same concentration. Staining for all antibodies was considered positive if more than 10% of the cells showed distinct reactivity. If more than one antibody was used for the same antigen, positive staining of one antibody is sufficient for positivity (for *p53* only, Do-1 or/and Do-7).

#### DNA isolation and PCR

The DNA from paraffin sections was isolated according to standard methods and PCR for exons 4 to 9 of the *p53* gene was performed as described (Taubert *et al.*, 1995). PCR for exons 1 and 2 of the *K-ras* gene (Grimmond *et al.*, 1992) and the *N-ras* gene (Svänen *et al.*, 1992) was performed as reported.

#### SSCP analysis and DNA sequencing

Non-radioactive SSCP analysis and DNA sequencing are described in detail (Thamm, 1995). Briefly: for SSCP analysis, 10 µl (approx. 1.5 µg) of PCR product were dissolved in SSCP buffer (98% formamide, 20 mM EDTA, 0.05% bromophenol blue), denatured for 5 min at 98°C and immediately stored on

ice. The samples were run in 6% or 10% non-denaturing ready-made TBE-gels (Novex, San Diego, CA) at 9 to 13°C for 2.5 to 3 hr (85–100 V). Afterwards, the gels were silver-stained according to standard protocols (silver-staining kit, Promega, Madison, WI) to detect shifts in the single-strand-DNA pattern. For sequencing, the purified PCR products were amplified by a cyclic PCR using the corresponding 5'-biotinylated primers, and sequencing products were verified by chemiluminescence (CPD-Star, Tropix, Bedford, MA).

#### Allele-specific oligonucleotide hybridization (ASO)

The PCR products from genomic DNA of the patients and the control were denatured by heat and immediately stored on ice. Samples and the control (5 µl each) were spotted on a nylon membrane, dried and UV-cross-linked for 5 min. After pre-hybridization (1 hr, 65°C) in 50 ml hybridization solution (10 × Denhardt's, 2 × SSC, 0.1% SDS), 100 ng 3'-biotin-labelled probe were added to 50 ml of freshly prepared hybridization solution (heat-denatured) and incubated overnight at 65°C. Washing of the membrane (15 min, 2 × SSC, 2 × 15 min, 1 × SSC) was performed at special temperatures (64°C, 68°C and 72°C). For the detection of DNA, a hybridization chemiluminescence assay (CPD-Star, Tropix) was applied.

Wild-type probe: E7wil24

5'-TCCGGTTCATGCCGCCATGCAGGG-3'

Mutant probe: E7mut24

5'-TCCGGTTCATGCTGCCATGCAGGG-3'

## RESULTS

#### Immunohistochemistry

Three groups of malignant STS — fibrosarcomas, malignant neural tumors (peripheral neuroblastoma and malignant PNST) and myogenic sarcomas (rhabdo- and leiomyosarcomas) — were investigated for *p53* and mdm2 protein, and their relationship to the grading was recorded (Table II). A total of 198 samples from 127 patients originated from 54 myogenic sarcomas (12 rhabdomyosarcomas and 42 leiomyosarcomas from 9 and 28 patients respectively), 63 malignant neural tumors (42 patients) and 81 fibrosarcomas (48 patients).

#### *p53* immunoreactivity

Of the tumor samples, 57% (113/198) showed immunohistochemically *p53*-positive after staining with DO-1 and/or DO-7 MAbs (Fig. 1 and 4). Separating the samples according to tumor entities, 75% of the rhabdomyosarcomas (9/12), 57% of the leiomyosarcomas (24/42), 54% of the malignant neural tumors (34/63) and 57% of the fibrosarcomas (46/81) showed *p53* positivity (Table II). Among these, 26% (4/15) grade-I, 58% (46/80) grade-II and 61% (63/103) grade-III samples were found; this reveals a correlation between increasing malignancy and the number of *p53*-positive tumors.

TABLE I - CHARACTERISTICS OF THE SOFT-TISSUE TUMOR SAMPLES<sup>1</sup>

Soft-tissue tumors	Myogenic sarcoma	Malignant neural tumors	Fibrosarcoma	Total
Tumor samples	54 <sup>2</sup> (35) <sup>3</sup>	63 <sup>4</sup> (31) <sup>5</sup>	81 (34)	198 (100) <sup>6</sup>
Primary tumors	30 (17)	38 (16)	40 (17)	108 (50)
Recurrences	15 (9)	17 (10)	38 (15)	70 (34)
Metastases	9 (9)	8 (5)	3 (2)	20 (16)
Grade 1	2 (2)	1 (1)	13 (6)	16 (9)
Grade 2	24 (19)	30 (14)	29 (18)	83 (51)
Grade 3	28 (14)	32 (16)	39 (10)	99 (40)
Patients	37 (26)	42 (25)	48 (27)	127 (78)
Patients alive	10 (6)	8 (3)	20 (8)	38 (17)
Patients dead	27 (20)	34 (22)	28 (19)	89 (61)

<sup>1</sup>Tumor samples and number of patients with soft-tissue tumors investigated immunohistochemically and (in parentheses) molecularly.—Includes 12 rhabdomyosarcoma (9 patients) and 42 leiomyosarcoma samples (28 patients). All but one of the rhabdomyosarcomas were adult, pleomorphic tumors.—Includes 6 rhabdomyosarcoma (3 patients) and 29 leiomyosarcoma samples (23 patients).—Consisting of 52 neurogenic sarcoma (33 patients) and 11 peripheral neuroblastoma samples (9 patients).—Consisting of 22 neurogenic sarcoma (20 patients) and 9 peripheral neuroblastoma samples (5 patients).—In most cases, a tumor is represented by one sample; a maximum of 3 samples originated from one tumor.

TABLE II - RESULTS OF IMMUNOHISTOCHEMICAL ANALYSIS FOR TUMOR SAMPLES OF THE TUMOR GROUPS MYOGENIC SARCOMAS, MALIGNANT NEURAL TUMORS AND FIBROSARCOMAS

STS samples positive / total	Myogenic sarcoma		Malignant neural tumors	Fibrosarcoma	Total
	Rhab	Leio			
p53 <sup>1</sup>	9/12	24/42	34/63	46/81	113/198
Grade 1	0/0	0/1	0/1	4/13	4/15
Grade 2	1/2	10/22	21/30	14/26	46/80
Grade 3	8/10	14/19	13/32	28/42	63/103
mdm2 (IF-2)	4/11	6/43	15/63	36/81	61/198
Grade 1	0/0	0/1	0/1	8/13	8/15
Grade 2	0/2	5/22	9/30	10/27	24/81
Grade 3	4/9	1/20	6/32	18/41	29/102

<sup>1</sup>Staining of p53 antibodies Do 1 and/or Do 7 was considered as positive.—Rhab, rhabdomyosarcoma; Leio, leiomyosarcoma.

*mdm2 immunoreactivity*

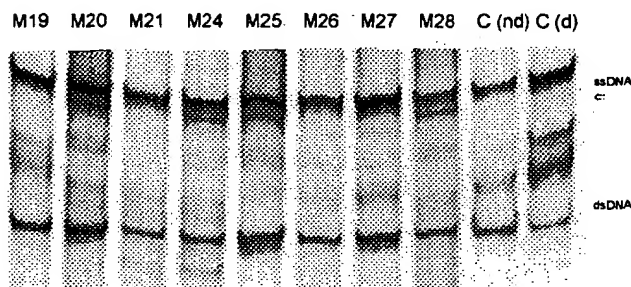
IF-2 was positive in 31% (61/198) of the tumor samples immunohistochemically studied. Positive staining was observed in 36% (4/11) of the rhabdomyosarcomas, 14% (6/43) of the leiomyosarcomas, 24% (15/63) of the MNT and 44% (36/81) of the fibrosarcomas (Table II, Fig. 1 and 6).

In contrast to the findings of Cordon-Cardo *et al.* (1994), mdm2 over-expression could not be related to a higher tumor grade, as shown also by Wiethge *et al.* (1994). However, over-expression of mdm2 protein occurs in all investigated soft-tissue entities, confirming the role of mdm2 over-expression in soft-tissue tumorigenesis (Leach *et al.*, 1993).

*Mutational analysis for p53 (exons 4 to 9)*

On investigation, 3 groups of soft-tissue sarcomas for *p53* mutations (exons 4 to 9), no such mutations could be identified for the MNT and fibrosarcoma entities (31 and 34 samples respectively), but mutations were detected in the group of myogenic sarcomas, *i.e.*, in leiomyosarcoma as well as in rhabdomyosarcoma samples.

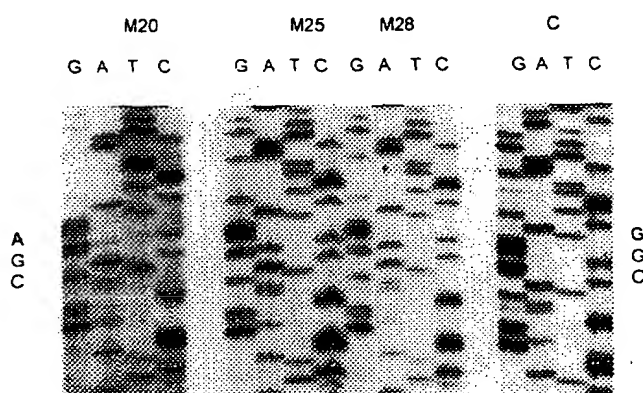
For 10/35 myogenic-sarcoma samples (5 from primary tumors, 4 from recurrences and one metastasis) from 6/26 patients, 4 different mutations were recorded. These mutations were G-to-A transitions in codon 158 and 245 respectively, a 1-bp insertion in codon 215 and a 15-bp deletion in exon 5 (Table III, Fig. 2). Surprisingly, 6 myogenic sarcoma samples from 3 patients carried the same mutation, a G-to-A transition in codon 245 (exon 7) (Table III). The samples originated from one recurrence (M28), 3 biopsies of a recurrence (M19, M20, M21) and 2 biopsies of a primary tumor (M24, M25) respectively. Samples M20 and M28 showed unambiguously the transition after sequencing. However, samples M19 and M21 (from the same tumor as M20) were not unambiguous, showing a very weak sequencing signal for the transitional base exchange. Consequently, allele-specific oligonucleotide hybridization (ASO) was applied, using mutant-specific (E7mut24) and wild-type-specific probes (E7wil24).



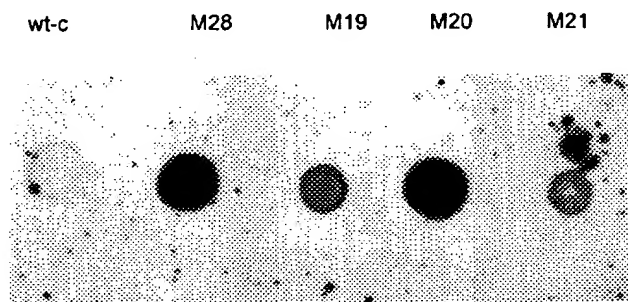
**FIGURE 1 – Result of SSCP analyses of exon 7 from the *p53* gene for myogenic sarcomas. A shift in the ssDNA pattern was observed in samples M20, M24, M25 and M28 (marked by an arrowhead). \*c-control, ssDNA, single-strand DNA; dsDNA, double-strand DNA. "nd-not denatured, d-denatured.**

The mutant-specific probe hybridized at 68°C only with potentially mutated DNA samples (M19/M20/M21, M28), but not with normal control DNA (Fig. 3). The wild-type-specific probe, on the other hand, hybridized with all DNA samples, because of wt-*p53*-alleles or remnants of normal cells (*e.g.*, infiltrating lymphocytes and vessels) still present in the sample (data not shown). The G-to-A transition was confirmed by repeating the ASO experiments, the PCR and sequencing reactions independently at least twice.

For one primary tumor of a leiomyosarcoma (M44) and its metastasis (M45) a G-to-A transition in codon 158 (exon 5)



**FIGURE 2 – Results of sequencing of myogenic sarcomas with a point mutation in codon 245. In all 3 tumor samples a GGC to AGC transition was identified in nucleotide 733.**



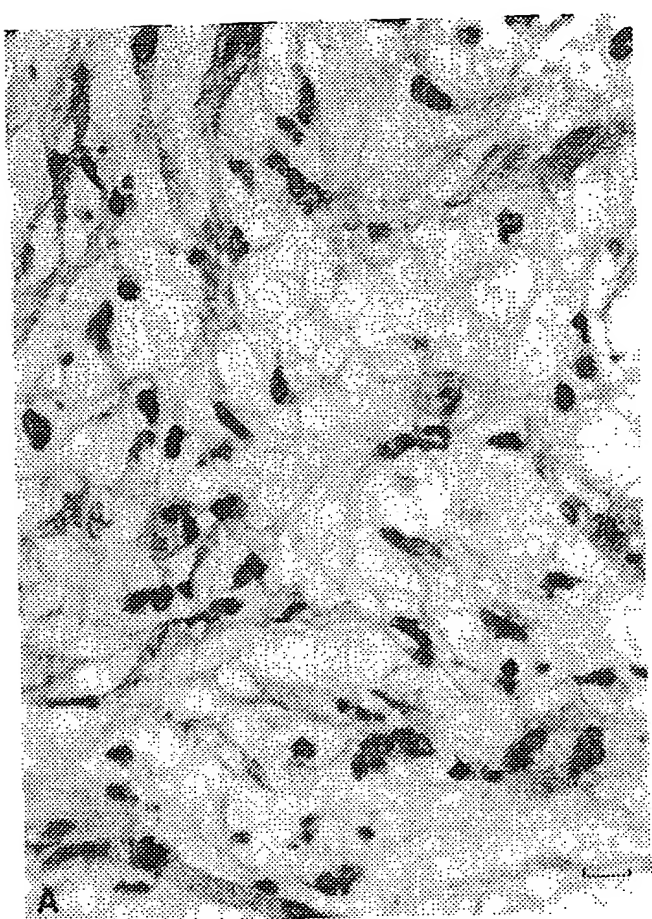
**FIGURE 3 – Results of the dot-blot of tumor samples from myogenic sarcomas (M28, M19, M20, M21) with wild-type-specific (wt-KE7) and mutant-type-specific oligonucleotides (mt-KE7) of PCR products for exon 7 of the *p53* gene. Hybridization with mutant-specific oligonucleotides (see "Material and Methods") at 68°C shows a specific signal for the tumor samples M28 (transition in codon 245 identified by sequencing) and M19, M20, M21. The control DNA (wt-cj from peripheral-blood cells) shows only a very weak signal.**

**TABLE III – RESULTS OF MOLECULAR ANALYSIS FOR MYOGENIC SARCOMAS WITH A *p53* MUTATION**

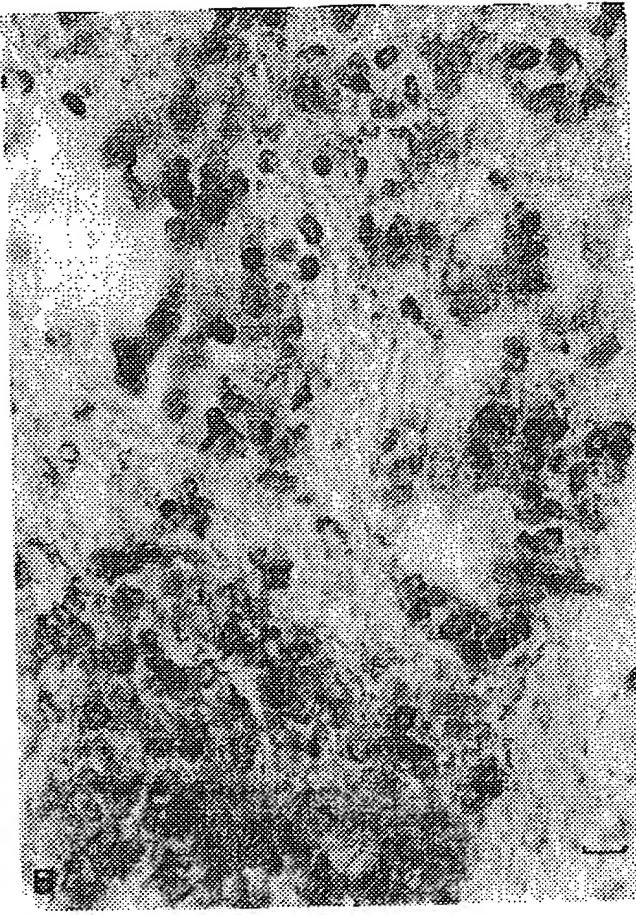
Tumor sample	Entity	Grade	P/R/M	sv	ex	nt	codon	alt	bp-alt	aa-alt
M42	Leio	III	P	d	4	318-332	106-111	del	-(15)	5-aa-del
M44/M45	Leio	II/II	P/M	d	5	473	158	ts	CGC → CAC	Arg → His
P6-93	Leio	III	P	a	6	643	215	ins	+ (1)	frameshift <sup>1</sup>
M24/M25	Leio	II/II	P/P	d	7	733	245	ts	GGC → AGC	Gly → Ser
M28	Rhab	II	R	d	7	733	245	ts	GGC → AGC	Gly → Ser
M19/M20/M21	Rhab	III/III/II	R/R/R	d	7	733	245	ts	GGC → AGC	Gly → Ser

All identified *p53* mutations are in the region of codons 106–245, and therefore concern the core protein domain (codons 100–300). The high portion of transitional-point mutations is striking. This was also observed for other soft-tissue tumor entities.

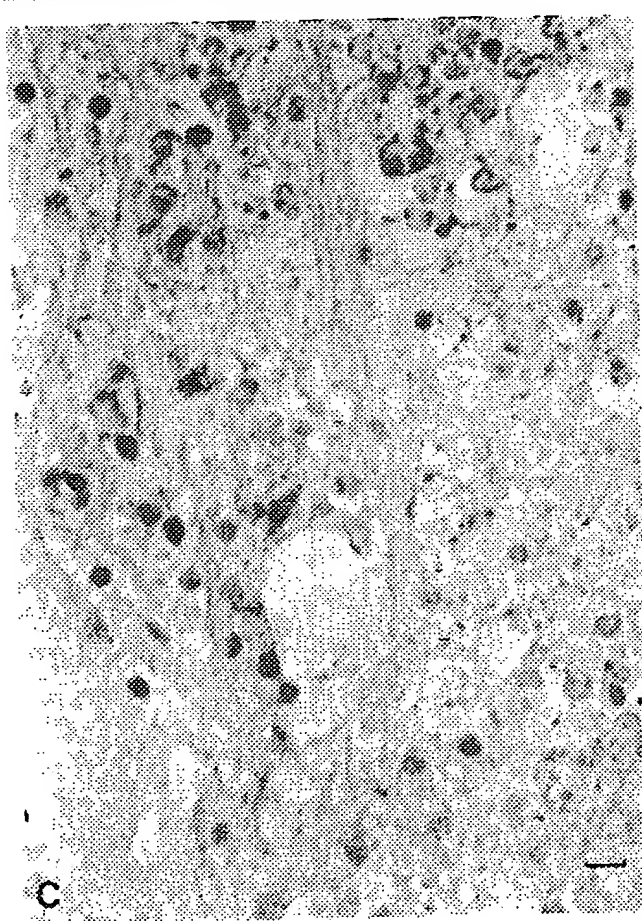
<sup>1</sup>The insertion identified results in a frameshift and a new stop codon (codon 221). P, primary tumor; R, recurrence; M, metastases; sv, survival; ex, exon; nt, nucleotide; bp, base pair(s); aa, amino acid(s); alt, alteration; del, deletion; a, alive; d, dead; Rhab, rhabdomyosarcoma; Leio, leiomyosarcoma; ts, transition; ins, insertion.



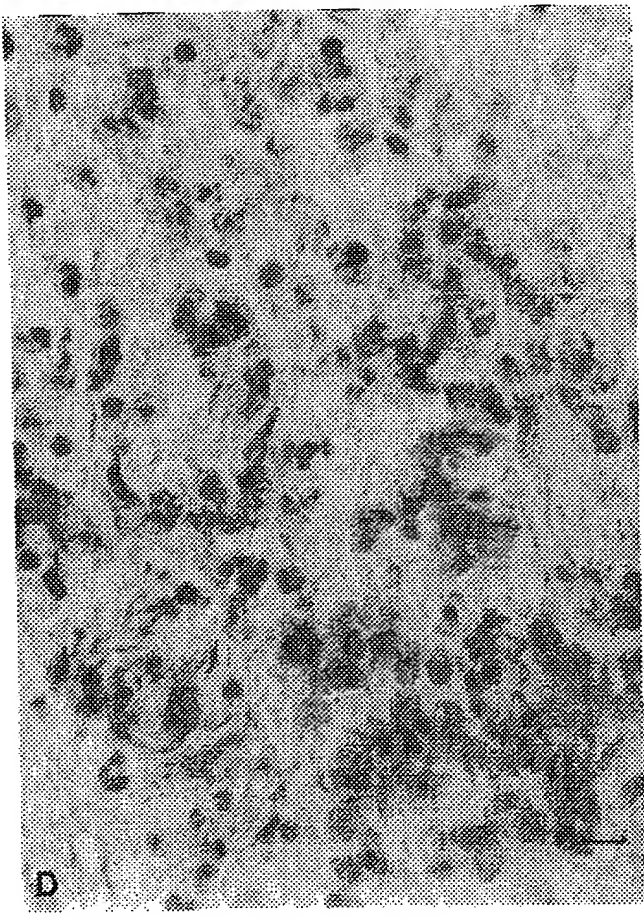
A



B



C



D

FIGURE 4

TABLE IV - RESULTS OF IMMUNOHISTOCHEMISTRY FOR p53 AND mdm2 IN MYOGENIC SARCOMA SAMPLES WITH p53 MUTATIONS

Sample	Grade	p53					mdm2 IF-2
		CM-1	DO-1	DO-7	Pab180I	Pab240	
M 19	III	+	-	-	-	+	-
M 20	III	+	++	++	+	++	++
M 21	II	++	-	-	-	++	-
M 24	II	++	++	++	++	++	-
M 25	II	++	++	++	++	+	-
M 28	II	++	++	++	++	+	-
M 42	III	+	+	-	-	+	-
M 44	II	-	++	++	+	+	-
M 45	II	+	++	++	+	-	+
P6-93	III	++	++	++	-	++	++

Assessment of applied antibodies: -, no expression; +, distinct expression; ++, strong expression.

was identified, pointing to a selective advantage of clones with this p53 mutation. Furthermore, a primary tumor of a leiomyosarcoma (P6-93) carried a 1-bp insertion in codon 215, causing a frame shift resulting in a stop in codon 221. This alteration was also detectable in a simultaneously established primary-cell culture of the same tumor (data not shown). Additionally, in one leiomyosarcoma sample (M42) a 15-bp deletion (codons 106–111) was detected, resulting in the loss of 5 amino acids and an amino-acid exchange from Ser to Arg in codon 106. In all sequencing reactions, the mutated sequence as well as the wt-sequence could be found as described above.

In a PCR-SSCP-sequencing analysis of exons 1 and 2 from the K-ras gene in myogenic sarcomas (32 tumor samples from 21 patients) and of exons 1 and 2 from the N-ras gene in MNT (31 tumor samples from 25 patients), no mutation could be detected (data not shown).

## DISCUSSION

Three groups of STS, i.e., fibrosarcomas, malignant neural tumors and myogenic sarcomas, were analyzed molecularly and immunohistochemically.

### Immunohistochemistry

The finding of 26% grade-I, 57% grade-II and 61% grade-III tumor samples with p53 positivity is comparable to other STS, such as MFH and liposarcoma, where grade-II and grade-III tumors in particular showed p53 positivity (Kawai *et al.*, 1994; Taubert *et al.*, 1995).

The result of 32% mdm2 positivity is similar to the findings of Cordon-Cardo *et al.* (1994), who found positive staining in 37% (76/207) of STS. Unfortunately, investigation for mdm2-gene amplification was not possible, since paraffin-embedded material was studied. However, the finding that all of the 33% (8/24) STS with mdm2-gene amplification also showed mdm2 over-expression (Leach *et al.*, 1993) suggests gene amplification as a possible reason for mdm2 over-expression.

We were able to support that mdm2 expression is more abundant in metastases than in primary tumors (Ladanyi *et al.*, 1993; Cordon-Cardo *et al.*, 1994), since mdm2 positivity was detected in M45 (metastasis), but not in M44 (primary tumor of the same leiomyosarcoma).

Generally, a combination of p53 and mdm2 over-expression is recorded for all STS entities in this study. Over-expression of mdm2 has been discussed mainly as an alternative to p53 alteration/over-expression of mt-p53 through inactivating p53

function (Leach *et al.*, 1993; Patterson *et al.*, 1994). However, cases of simultaneous over-expression for p53 and mdm2 have also been recorded (Cordon-Cardo *et al.*, 1994; Marston *et al.*, 1994). It is suggested that p53 protein over-expression may induce increased mdm2 RNA transcription (Florenes *et al.*, 1994), which could result in over-expression of mdm2 protein. Moreover, p53-mdm2 complexes could activate a function promoting tumorigenesis (Landers *et al.*, 1994).

Expression of mdm2 in p53-mutated myogenic sarcomas shows a heterogeneous picture. A 15-bp deletion (M42) showed no mdm2 positivity. An 1-bp insertional mutation was positive for mdm2, and the tumor samples with transitional mutations were in part mdm2-positive (Table IV). However, the latter result depended on the amount of tumor material (comparable to results in the sequencing reactions): for example, the M20 sample expressed strong mdm2 positivity, whereas M19 showed none. But at least one tumor sample (M20, M21, M25, M28, M45) from a patient with G-to-A transitions always showed mdm2 expression also. The co-existence of p53 mutations and mdm2 positivity could be explained by a selective advantage in tumors of weaker phenotype (Ladanyi *et al.*, 1993).

### Mutational analysis

Molecular characterization comprised a PCR-SSCP-sequencing analysis for the tumor-suppressor gene p53 (exons 4 to 9) and in part for the N-ras and K-ras genes (exons 1 and 2).

Neither N-ras nor K-ras mutations could be detected. This agrees with other studies, which found no K-ras mutations (Pulciani *et al.*, 1982; Wilke and Robinson, 1993) and just 3 N-ras mutations in STS. All these mutations concerned codon 61 of exon 2, identified in a neuroblastoma, a fibrosarcoma and a rhabdomyosarcoma (Taparowsky *et al.*, 1983; Brown *et al.*, 1984; Chardin *et al.*, 1985). However, N-ras- and K-ras-gene mutations do not seem to play an important role in STS tumorigenesis.

p53 mutational analysis revealed 6 mutations in 35 myogenic sarcoma samples, but none in 34 fibrosarcoma and 31 MNT samples. In previous mutational studies, p53 mutations appeared somewhat rarely in fibrosarcoma and MNT. The only p53 mutation described in a fibrosarcoma (Latres *et al.*, 1994) represents an exceptional occurrence of p53 mutations in this STS entity. In MNT, a small percentage may carry p53 mutations, as shown for 4 cases of neuroblastomas (Imamura *et al.*, 1993; Vogan *et al.*, 1993; Hosoi *et al.*, 1994) and 2 cases of neurofibrosarcomas (Menon *et al.*, 1990). But mutational frequencies in the range of 11 to 18% verified for other soft-tissue entities (Stratton *et al.*, 1990, 10/94; Toguchida *et al.*, 1992, 3/17; Leach *et al.*, 1993, 4/24; Taubert *et al.*, 1995), were not recorded for fibrosarcomas and MNT.

For 10/35 tumor samples (6/29 leiomyosarcomas and 4/6 rhabdomyosarcoma samples) 6 mutations were identified. This

FIGURE 4 – Immunohistochemical staining of rhabdomyosarcoma M20 with anti-p53 and anti-mdm2 antibodies. (a) hemalaun/eosin staining; (b) anti-mdm2 antibody IF-2; (c) anti-p53 antibody DO-1; (d) anti-p53 antibody DO-7. Scale bars, 10 µm.

result is comparable to the finding of 7 mutations in 26 myogenic sarcoma samples (4 mutations in 20 leiomyosarcomas and 2 mutations in 6 rhabdomyosarcomas) by Stratton *et al.* (1990). Unfortunately, no results concerning the patients are presented in this study. However, we found that 23% (6/26) of the myogenic sarcoma patients (4/23 of leiomyosarcoma patients and 2/3 of rhabdomyosarcoma patients) carried *p53* mutations.

In exon 4 we identified a 15-bp deletion (M42), in exon 5 a CGC-to-CAC transition for codon 158 (M44/M45), in exon 6 a 1-bp insertion in codon 215 and, surprisingly, in exon 7 a GGC-to-AGC transition in codon 245, which was identical in 3 patients (M19-21; M24/M25; M28). All mutations are located inside the core domain (codons 102-292; Cho *et al.*, 1994) and seem to affect structural rather than functional properties of the *p53*-DNA interaction.

All identified point mutations are G-to-A transitions and occur at CpG dinucleotides. It is known that CpGs are preferential loci for mutational hot spots. Although CpG sequences are under-represented in the human genome by the factor of 5, about 35% of point mutations causing human disorders occur at CpGs, and over 90% of these are transitions from G-to-A (Cooper and Youssoufian, 1988). In addition to causes such as differences in the fidelity and strand specificity of eucaryotic polymerases (Kunkel and Alexander, 1985; Wu and Maeda, 1987), cytosine methylation at CpG sites may cause the high mutational frequency. 5-methylcytosine is attackable by de-amination, whereby the [5-methyl]-cytosine is replaced by a thymine residue (a guanine by an adenine residue on the other strand; *i.e.*, a G-to-A transition) (Coulondre *et al.*, 1978; Lindahl and Nyberg, 1974). The resulting T:G mismatches cause minor distortions of the DNA helix (Brown

*et al.*, 1985) and it is more difficult for repair enzymes to recognize them. T:G mismatches are by a factor of 6000 less efficiently repaired than U:G mismatches formed by de-amination of cytosine (Schmutte *et al.*, 1995).

What is remarkable about the *p53* gene is that 5 out of 6 *p53* mutation hot-spot codons contain CpG dinucleotides (175, 245, 248, 273 and 282). This implies methylation-driven de-amination of 5-methyl cytosine as a major source of G-to-A transition mutations at CpG dinucleotides (Tornaletti and Pfeifer, 1995). The CpG site at codon 245 is well characterized as a mutational hot spot in carcinomas, with a total of 144 mutational cases out of 4496 entries (3.2%) in the *p53* mutation databank (Hollstein *et al.*, 1996). Of these, 67 cases concern the G-to-A transition. In sarcomas, no mutational hot spot has been described (Greenblatt *et al.*, 1994). For codon 245, 5 mutation cases out of 162 mutation entries for sarcomas are compiled (Hollstein *et al.*, 1996); if we add the 3 described here, the total recorded cases number 8 (*i.e.*, 5% of known sarcoma mutations). Of these, 6 cases have a G-to-A transition. Summarizing results, codon 245 appears as a mutational hot spot for sarcomas.

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Set	Items	Description
S1	2927712	CANCER OR TUMOR OR MALIGNANT?
S2	135446	P5 <sup>3</sup>
S3	101820	S1 AND S2
S4	2203	MDM2 (5N)EXPRESS?
S5	1672	S3 AND S4
S6	187	S5 AND PY<=1996
S7	1129	(AMPLIFICATION OR OVEREXPRESS?) (5N) MDM2
S8	114	S6 NOT S7
S9	66	RD (unique items)
? s increases?		
Sending Break...		
?s increases? (5n) mdm2		
5899392 INCREASES?		
7904 MDM2		
S10 623 INCREASES? (5N) MDM2		
? s s9 not s10		
66 S9		
623 S10		
S11 60 S9 NOT S10		
? t s11/3,k,ab/1-20		

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11718537 PMID: 9275659  
**[Studies on MDM2 oncogene expression and its effect on pancreatic carcinoma cells]**  
 Guo H; Liu T; Gao J  
 Department of Pathology, PUMC Hospital, CAMS, Beijing.  
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Over-expression of the MDM2 gene is found in some cases of haematological malignancies .

Quesnel B; Preudhomme C; Oscier D; Lepelley P; Collyn-d'Hooghe M; Facon T

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We looked for **MDM2** gene amplification and over- **expression** by Southern

and Northern blot analysis in 135 and 66 cases of haematological

**malignancies** , including ALL, AML, CML in chronic phase, CLL, MDS, PLL,

non-Hodgkin's lymphoma (NHL) and myeloma. No amplification of the gene was

found. An over- **expression** of **MDM2** RNA was seen in 9/66 (14%) patients

tested, including 3/9 ALL, 3/24 AML, 2/4 myelomas, 1/1 PLL, but 0/2 CML,

0/2 NHL and 0/21 MDS. None of the patients over- **expressing** **MDM2** had

modifications of **P53** gene transcript or **p53** mutations. Most of the

patients over- **expressing** **MDM2** gene had poor prognostic features

(including 'unfavourable' cytogenetic abnormalities), poor response to

chemotherapy and short survival. Our findings suggest that over- **expression**

of **MDM2** is seen in a relatively small number of haematological

**malignancies** , and is associated with poor prognosis.

Over-expression of the MDM2 gene is found in some cases of haematological malignancies .

Oct 1994 ,

We looked for **MDM2** gene amplification and over- **expression** by Southern

and Northern blot analysis in 135 and 66 cases of

haematological  
**malignancies** , including ALL, AML, CML in chronic phase, CLL,  
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non-Hodgkin's lymphoma (NHL) and myeloma. No amplification of  
the gene was  
found. An over- **expression** of **MDM2** RNA was seen in 9/66  
(14%) patients  
tested, including 3/9 ALL, 3/24...

...0/2 CML, 0/2 NHL and 0/21 MDS. None of the patients over-  
**expressing**

**MDM2** had modifications of **P53** gene transcript or **p53**  
mutations. Most

**Amplification of the MDM2 gene in human breast cancer  
and its  
association with MDM2 and p53 protein status.**

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The present study reports on the frequency of **MDM2** gene amplification

and **MDM2** protein **expression** in a series of 100 breast carcinomas and

its association with accumulation of the **p53** protein. Of the 100 cases,

frozen samples for 82 cases were available for Southern blotting. Three of

the 82 (4%) demonstrated MDM2 gene amplification of up to 6-fold.

Immunohistochemical analysis of the formalin-fixed, paraffin-embedded

tumours demonstrated that 7/97 (7%) had nuclear **expression** for **MDM2** in

10-50% of the tumour cells (type 2 staining) and were denoted MDM2+. Two of

the MDM2-amplified samples were MDM2+ with one of the two tumours also

displaying type 2 **p53** nuclear staining. Finally at the protein level,

MDM2+ tumours were significantly associated with tumours having low levels

of **p53** staining (0-10% cells positive) ( $P = 0.03$ ). We conclude that MDM2

gene amplification occurs at a lower frequency in breast **cancer** than in

non-epithelial tumours. Alterations in MDM2 and **p53** may represent

alternative pathways in tumorigenesis, but they are not mutually exclusive in all cases.

Frequent occurrence of p53 mutations in rhabdomyosarcoma and leiomyosarcoma, but not in fibrosarcoma and malignant neural tumors.

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We have analyzed soft-tissue sarcomas (STS) molecularly for mutations in

the **tumor** -suppressor gene **p53** and immunohisto- chemically for

**expression** of **p53** and **mdm2** proteins. In this study, **tumor** samples

from 3 groups of soft-tissue sarcomas, i.e., fibrosarcomas, myogenic

sarcomas and **malignant** neural tumors (MNT), were investigated. The

methods applied encompass immunohistochemistry on 198 **tumor** samples using

**p53** antibodies (DO-1 and DO-7) and an mdm2 antibody (IF-2). Out of these, 100 samples were subjected to non-radioactive PCR-SSCP-sequencing analysis.

Immunohistochemical detection rate for **p53** (range of 57% to 67%) and for

mdm2 proteins (range of 19 to 44%) was similar in all 3 groups. In higher

**tumor** grades, an increased rate of immunopositivity was found for **p53**

but not for mdm2. Investigation of **p53** mutational status revealed 6

mutations in myogenic sarcomas but none in **malignant** neural tumors or

fibrosarcomas, suggesting different roles of **p53** in the 3 STS groups.

Interestingly, a G-->A transition in codon 245 (a CpG site) was found in 3

myogenic sarcomas. Our results and those of others suggest **p53** codon 245 as a mutational hotspot in sarcomas, as recognized in carcinomas.

**Frequent occurrence of p53 mutations in rhabdomyosarcoma and leiomyosarcoma, but not in fibrosarcoma and malignant neural tumors.**

Aug 22 1996 ,  
We have analyzed soft-tissue sarcomas (STS) molecularly for mutations in

the **tumor** -suppressor gene **p53** and immunohisto- chemically for **expression** of **p53** and **mdm2** proteins. In this study, **tumor** samples

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methods applied encompass immunohistochemistry on 198 **tumor** samples using

**p53** antibodies (DO-1 and DO-7) and an mdm2 antibody (IF-2). Out of these, 100 samples were subjected to non-radioactive PCR-SSCP-sequencing analysis.

Immunohistochemical detection rate for **p53** (range of 57% to 67%) and for **mdm2** proteins (range of 19 to 44%) was similar in all 3 groups. In higher

**tumor** grades, an increased rate of immunopositivity was found for **p53** but not for **mdm2**. Investigation of **p53** mutational status revealed 6 mutations in myogenic sarcomas but none in **malignant** neural tumors or fibrosarcomas, suggesting different roles of **p53** in the 3 STS groups.

Interestingly, a G-->A trans

Set	Items	Description
S1	2927712	CANCER OR TUMOR OR MALIGNAN?
S2	135446	P53
S3	101820	S1 AND S2
S4	2203	MDM2 (5N) EXPRESS?
S5	1672	S3 AND S4
S6	187	S5 AND PY<=1996
S7	1129	(AMPLIFICATION OR OVEREXPRESS?) (5N) MDM2
S8	114	S6 NOT S7
S9	66	RD (unique items)
? s increas?		
Sending Break...		
?s increas?	(5n)mdm2	
5899392	INCREAS?	
7904	MDM2	
S10	623	INCREAS? (5N) MDM2
? s s9 not s10		
	66	S9
	623	S10
S11	60	S9 NOT S10
? t s11/3, k, ab/1-20		

**APOPTOSIS, CANCER AND THE P53 TUMOR-SUPPRESSOR GENE** (Abstract Available)

Author(s): LEE-JM; BERNSTEIN A

Corporate Source: MT SINAI HOSP, SAMUEL LUNENFELD RES INST, DIV MOLEC & DEV

BIOL, 600 UNIV AVE/TORONTO/ON M5G 1X5/CANADA/; MT SINAI HOSP, SAMUEL

LUNENFELD RES INST, DIV MOLEC & DEV BIOL/TORONTO/ON M5G 1X5/CANADA/;

UNIV TORONTO, DEPT MOLEC & MED GENET/TORONTO/ON M5S 1A8/CANADA/

Journal: CANCER AND METASTASIS REVIEWS, 1995 , V14, N2 (JUN), P149-161

ISSN: 0167-7659

Language: ENGLISH Document Type: REVIEW

Abstract: One of the most commonly detected abnormalities in human cancer

is mutation of the p53 tumour suppressor gene. Intrinsic to the

function of p53 is its ability to induce apoptotic cell death and to

cause **cell cycle arrest**. Moreover, p53 plays an important role in

controlling the cellular response to DNA damaging agents such as

ionizing radiation and cancer chemotherapeutic drugs. Loss of p53

function causes increased resistance to radiation and chemotherapeutic

agents, and there is increasing evidence that p53 mutational status is

an important determinant of clinical outcome in cancer. This review

will focus on recent data describing the biochemistry of p53 function,

its role in mediating apoptosis and **cell cycle arrest** and in the

control of tumour growth and death.

, 1995

**MARKER GENES FOR CYTOTOXIC EXPOSURE - P53 (Abstract Available)**

Author(s): MONTENARH M

Corporate Source: UNIV SAARLAND, BLDG 44/D-66424 HOMBURG//GERMANY/

Journal: STEM CELLS, 1995 , V13, S1 (MAY), P136-141

ISSN: 1066-5099

Language: ENGLISH Document Type: ARTICLE

Abstract: The growth suppressor p53 plays an important role in the

regulation of cell proliferation, DNA repair and apoptosis.

In

wild-type p53 expressing cells, gamma-irradiation induces an increase

in the level of p53 protein and these cells exhibit a GI growth arrest.

The p53-induced G(1) growth arrest is abrogated in cells expressing

mutant p53, or in cells where p53 is inactivated by complex formation

with cellular or viral proteins such as mdm2 or the E6 proteins of

human papillomavirus (HPV) 16 or HPV18. Wild-type p53 expressing cells

are radiosensitive whereas mutant p53 expressing cells are radioresistant. In some cell types, p53 mutations are observed after

gamma-irradiation of cells although this observation is not consistent

for all cell types. Furthermore, it is not clear whether these

mutations are the direct result of irradiation or secondary effects.

, 1995

```
? s p53
    S1 135504 P53
? s activat? (5n) p53
    2455436 ACTIVAT?
    135504 P53
    S2 11161 ACTIVAT? (5N) P53
? s cell(w)cycle(w)arrest
    5633738 CELL
    750341 CYCLE
    142979 ARREST
    S3 21150 CELL(W)CYCLE(W)ARREST
? s s2 and s3
    11161 S2
    21150 S3
    S4 1578 S2 AND S3
? s transcription?(5n)factor
    861751 TRANSCRIPTION?
    2373148 FACTOR
    S5 216100 TRANSCRIPTION?(5N)FACTOR
? s s4 and s5
    1578 S4
    216100 S5
    S6 333 S4 AND S5
? s s6 and py<1997
Processing
    333 S6
    31263329 PY<1997
    S7 77 S6 AND PY<1997
? rd
>>>Duplicate detection is not supported for File 340.

>>>Records from unsupported files will be retained in the RD set.
...examined 50 records (50)
...completed examining records
    S8 72 RD (unique items)
? t s8/3,k,ab/60-72
```

**8/3,K,AB/60 (Item 53 from file: 34)**  
DIALOG(R)File 34:SciSearch(R) Cited Ref Sci  
(c) 2005 Inst for Sci Info. All rts. reserv.

04235094 Genuine Article#: RQ469 Number of References: 63  
**Title: P53 DEPENDENT GROWTH SUP**

```
? s cancer? or tumor or malignan?
    1585336  CANCER?
    1597891  TUMOR
    589815   MALIGNAN?
    S1 2963795 CANCER? OR TUMOR OR MALIGNAN?
? s p53
    S2 135446  P53
? s s1 and s2
    2963795  S1
    135446   S2
    S3 102760  S1 AND S2
? s mdm2
    S4 7904  MDM2
? s s3 and s4
    102760  S3
    7904   S4
    S5 5534  S3 AND S4
? s not(w)overexpress?
>>>Operator "NOT" in invalid position
? s overexpress?
    S6 209394 OVEREXPRESS?
? s s5 and s6
    5534  S5
    209394 S6
    S7 1726  S5 AND S6
? s s5 not s6
    5534  S5
    209394 S6
    S8 3808  S5 NOT S6
? s mdm2(5n) express?
    7904  MDM2
    3414915 EXPRESS?
    S9 2203  MDM2(5N) EXPRESS?
? s s3 and s9
    102760  S3
    2203   S9
    S10 1685  S3 AND S9
? s s10 and py<=1997
Processing
Sending Break...
?s s10 and py<1997
Processing
    1685  S10
    31263327 PY<1997
    S11 188   S10 AND PY<1997
? rd
>>>Duplicate detection is not supported for File 340.

>>>Records from unsupported files will be retained in the RD set.
...examined 50 records (50)
```

...examined 50 records (100)  
...examined 50 records (150)  
...completed examining records  
    S12     106 RD (unique items)  
? s excess  
    S13 328150 EXCESS  
? s s12 not s13  
    106 S12  
    328150 S13  
    S14     105 S12 NOT S13  
? s sarcoma  
    S15 147075 SARCOMA  
? s s14 not s15  
    105 S14  
    147075 S15  
    S16     101 S14 NOT S15  
? t s16/3,k,ab/1-5

**16/3,K,AB/1 (Item 1 from file: 155)**

DIALOG(R) File 155: MEDLINE(R)  
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11736055 PMID: 9815889  
**Differential expression of multiple MDM2 messenger RNAs and proteins in normal and tumorigenic breast epithelial cells.**  
Gudas J M; Nguyen H; Klein R C; Katayose D; Seth P; Cowan K H  
Medicine Branch, Division of Cancer Treatment, Medical  
Breast Cancer  
Section, National Cancer Institute, Bethesda, Maryland 20892,  
USA.  
Clinical cancer research - an official journal of the  
American  
Association for Cancer Research (UNITED STATES) Jan 1995  
, 1 (1)  
p71-80, ISSN 1078-0432 Journal Code: 9502500  
Publishing Model Print  
Document type: Journal Article  
Languages: ENGLISH  
Main Citation Owner: NLM  
Record type: MEDLINE; Completed  
The MDM2 gene is a nuclear phosphoprotein that is regulated by  
**p53** and  
functions, in one capacity, to inhibit the transcriptional  
activity of the  
wild-type **p53** protein. Multiple MDM2 transcripts were  
detected in human  
breast epithelial cells. In estrogen receptor-negative normal,  
immortal,  
and tumorigenic breast epithelial cells, we found a good

correlation between **MDM2** mRNA levels and **expression** of wild-type **p53**. When wild-type **p53** was overexpressed in estrogen receptor-negative **tumor** cells containing a mutant or no endogenous **p53**, MDM2 mRNA levels increased significantly, indicating that wild-type **p53** positively influences MDM2 mRNA levels in these **tumor** cells. Because all estrogen receptor-positive breast **tumor** cells had high MDM2 mRNA levels regardless of the status of their endogenous **p53** protein, other factors likely influence **MDM2 expression**.

[Studies on MDM2 oncogene expression and its effect on  
pancreatic  
carcinoma cells]

Guo H; Liu T; Gao J  
Department of Pathology, PUMC Hospital, CAMS, Beijing.  
Zhonghua bing li xue za zhi Chinese journal of pathology  
(CHINA) Aug  
**1996**, 25 (4) p232-5, ISSN 0529-5807 Journal Code: 0005331  
Publishing Model Print  
Document type: Journal Article ; English Abstract  
Languages: CHINESE  
Main Citation Owner: NLM  
Record type: MEDLINE; Completed  
In order to study the interrelation and interaction between  
MDM2 oncogene  
and wild type **p53** in human pancreatic **cancer** , we  
studied the  
**expression** and amplification of **MDM2** oncogene and its  
antagonistic  
effect on wild type **p53** by use of gene recombination, gene  
transduction  
and molecular hybridization techniques. The results showed  
that MDM2  
oncogene could be detected in all 5 pancreatic cell lines, but  
**MDM2** mRNA  
**expression** varied in the different cell lines. The  
recombinant vector  
pCMV-MDM2 was transduced into PC-2/s-wtp53 cell line (a  
transformed PC-2  
pancreatic carcinoma cell line containing wild type **p53**  
gene). The  
resultant cell line, PC-2/s-wtp53/pCMV-MDM2 showed rapid cell  
growth, a  
rate similar to that of the parent cell line PC-2. Our results  
verify the  
fact that MDM2 gene can abrogate the cell growth arrest  
mediated by wild  
type **p53** and the antagonistic function of wild type **p53** .

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